PTO/SB/08B (08-03)

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Substitute for form 1449B/PTO

## INFORMATION DISCLOSURE STATEMENT BY APPLICANT

(use as many sheets as necessary)

Sheet 3 of 3

Complete if Known				
Application Number	08/856,376			
Filing Date	May 14, 1997			
First Named Inventor	Chee, Mark			
Art Unit	1631			
Examiner Name	Ardin Marschel			
Attorney Docket Number	018547-025010US			

	NON PATENT LITERATURE DOCUMENTS						
Examiner Initials *	Cite No. <sup>1</sup>	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.	T²				
DM	AQ	Ruvolo et al., Mitochondrial COII sequences and modern huiman origins," Molecular Biology and Evolution, 10:1115 (1993).					
	AR	Seneca et al., "Importance of sequence analysis in the NARP syndrome," J. Inherited Metabolic Disorders, 18 (1):97 (1995).					
7	AS	Tanaka and Ozawa, "Strand asymmetry in human mitochondrial DNA mutations," Genomics, 22(2):327 (1994).					

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Examiner Signature	Adm	Monsoly	Date Considered	1-22-04

EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to applicant.

Applicant's unique citation designation number (optional). Applicant is to place a check mark here if English language Translation is attached.

PTO/SB/08A (08-03)

Substitut	te for form 1449A/P	TOG.	2	Complete if Known			
		B	ADEMARK OF	Application Number	08/856,376		
INFO	DRMATIO	N DI	SCLOSURE	Filing Date	May 14, 1997		
STA	STATEMENT BY APPLICANT			First Named Inventor	Chee, Mark		
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Sheet	1	of	3	Attorney Docket Number	018547-025010US		

	U.S. PATENT DOCUMENTS+						
Examiner Initials*	Cite No.	Document Number  Number Kind Code <sup>2</sup> ( <i>if known</i> )	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear		

	FOREIGN PATENT DOCUMENTS							
	0.11-	Forei	gn Patent Doo	cument		Name of Patentee or	Pages, Columns, Lines, Where Relevant	
Examiner Initials*	Cite No. <sup>1</sup>	Country Code <sup>3</sup>	Number <sup>4</sup>	Kind Code <sup>6</sup> ( <i>il known</i> )	Publication Date MM-DD-YYYY	Applicant of Cited Document	Passages or Relevant Figures Appear	T⁰

Examiner Signature Date Considered 1-22-04

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of 3 2 Sheet

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AM	AA	Ginther et al., "Identifying individuals by sequencing mitochondrial DNA from teeth," Nature Genetics, 2:135 (10/1992).	
	АВ	Greenberg et al., "Intraspecific nucleotide sequence variability surrounding the origin of replication in human mitochondrial DNA," Gene 21(1-2):33 (1983).	
	AC	Howell et al., "Mitochondrial gene segregation in humans: is the bottleneck always narrow?" Human Genetics, 90:117 (1992).	
	AD	Howell et al., "When does bilateral optic atrophy become Leber hereditary optic atrophy?" American Journal of Human Genetics, 53:959 (1993).	
	AE	Hutchin et al., "A molecular basis for human hypersensitivity to aminoglycoside antibiotics," NAR 21(18):4174 (1993).	
	AF	Ikebe et al., "Point mutations of mitochondrial genome in PArkinson's disease," Molecular Brain Research 28(2):281 (1995).	
	AG	Isenberg and Moore, "Mitochondrial DNA Analysis at the FBI Laboratory," Forensic Science Communications, Vol. 1, No. 2 (7/1999).	
	АН	Johns and Neufeld, "Pitfalls in the molecular genetic diagnosis of Leber hereditary optic neuropathy (LHON)," American Journal of Human Genetics, 53 (4):916 (1993).	
	Al	Marzuki et al., "Normal variants of human mitochondrial DNA and translation products: building a reference data base," Human Genetics, 88 (2):139 (1991).	
	AJ	Mehta, et al., "A new genetic polymorphism in the 16S ribosomal RNA gene of human mitochondrial DNA," Annals of Human Genetics, 53 (Pt. 4):303 (1989).	
	AK	Moraes, et al., "Two novel pathogenic mitochondrial DNA mutations affecting organelle number and protein synthesis. Is the tRNA Leu(UUR) gene an etiologic hot spot?" J. of Clinical Investigation, 92(6):2906 (1993).	
	AL	Ozawa et al., "Distinct clustering of point mutations in mitochondrial DNA amoung patients with mitochondrial encephalomyopathies and with Parkinson's disease," BBRC, 176 (2):938 (1991).	
	АМ	Ozawa et al., "Patients with idiopathic cardiomyopathy belong to the same mitochondrial gene family of Parkinson's disease and mitochondrial encephalomyopathy," BBRC 177(1):518 (1991).	
	AN	Petruzzella et al., "Is a point mutation in the mitochondrial ND2 gene associated with Alzheimer's disease?" BBRC 186:491 (1992).	
	AO	Prezant et al., "Mitochondrial ribosomat RNA mutation associated with both antibiotic-induced and non-syndromic deafness," Nature Genetics, 4 (3):289.	
	AP	Reid et al., "Complete mtDNA sequence of a patient in a maternal pedigree with sensorineural deafness," Human Molecular Genetics, 3(8):1435 (1994).	

Examiner Signature	[ \	In Marsh	2	Date Considered	1-22-04	

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